[19]

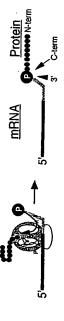


Fig. 1. Formation of mRNA-protein fusions on the ribosome. The ribosome pauses at an RNA/DNA junction, allowing puromycin to thread into the ribosome, entering the A site and forming fusion by accepting the nascent peptide from the peptidyl-tRNA in the P site.

reading and amplifying a protein sequence after it has been purified based library size that may be generated is limited by the size and efficiency of the translation reaction and by the efficiency of fusion formation on the ribosome. At the present time, libraries containing more than 1013 different sized as fusions commonly retain the binding properties of the unfused united in a single molecule, RNA-protein fusions provide a means for on its function, effectively reverse translating the protein. The maximum sequences can be readily generated. Further, peptides and proteins synthepolypeptides. Finally, proteins ranging in size from 1 to at least 30 kDa can be synthesized as fusions, thus opening a great diversity of systems to examination

Selection Scheme

The basic scheme in an RNA-protein fusion selection experiment is highlighted in Fig. 2, divided into 10 discrete steps: (1) generation of the

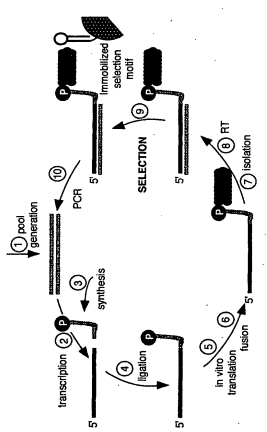


Fig. 2. Generalized selection scheme using mRNA-protein fusions. See text for description and optimization of individual steps.

KNA-PROTEIN FUSIONS

fusion, (7) isolation of the fusion, (8) reverse transcription to generate the immobilized selection motif, and (10) polymerase chain reaction (PCR) to igation of the puromycin oligonucleotide to the mRNA, (5) in vitro translaion of mRNA-puromycin templates, (6) generation of the mRNA-protein nitial double-stranded DNA sequence or pool, (2) transcription of the DNA into mRNA, (3) synthesis of the 3'-puromycin oligonucleotide, (4) cDNA/mRNA-protein fusion, (9) isolation of functional fusions with generate an enriched dsDNA pool.

1. dsDNA Library

lated region (5'UTR) should be chosen according to the in vitro translation system to be used for fusion generation. For translation in reticulocyte lysate, we commonly use a deletion mutant of the tobacco mosaic virus 5' tion in Escherichia coli, a Shine-Dalgarno sequence appropriately spaced A T7 promoter is present at the 5' end to allow large-scale synthesis of mRNA in vitro using T7 polymerase.13 The transcript begins with 3 G nucleotides to aid transcription initiation. The remainder of the 5'-untrans-JTR (ATMV) that provides efficient translation initiation. 12,14 For transla-The starting library is constructed as a mixture of double-stranded DNA sequences. The DNA sequence contains several important design features. with respect to the start codon should be chosen. 15,16

in reticulocyte lysate many sequences function efficiently as the 5' UTR.18 with a preference for A as the first purine (-3) and G as the second (+4). 1820 In contrast with bacteria, the 5' UTR in eukaryotes does not contain reports have been made of sequences that greatly enhance translation,17 In general, eukaryotic translation systems use the first AUG codon in the mRNA to initiate protein synthesis. The precise sequence context surrounding this codon influences the efficiency of translation. 18,19 The sequence 5'RNNAUGR provides a good start context for most sequences, a ribosome-binding site or an extensive 5' consensus sequence. Although

sequence(s) or a random sequence library. The most important feature of The open reading frame (ORF) can be constructed from either a defined he ORF and adjacent 3' constant region is that neither contain stop codons.

F. Milligan and O. C. Uhlenbeck, Methods Enzymol. 180, 51 (1989)

¹⁴ D. R. Gallie, D. E. Sleat, J. W. Watts, P. C. Turner, and T. M. A. Wilson, Nucleic Acid Res. 16, 883 (1988)

¹⁶ G. D. Stormo, T. D. Schneider, and L. M. Gold, Nucleic Acids Res. 10, 2971 (1982). 15 J. A. Steitz and K. Jakes, Proc. Natl. Acad. Sci. U.S.A. 72, 4734 (1975).

¹⁷S. A. Jobling and L. Gehrke, Nature 325, 622 (1987).

¹⁹ M. Kozak, J. Biol. Chem. 266, 19867 (1991). ¹⁸ M. Kozak, Microbiol. Rev. 47, 1 (1983)

²⁰ M. Kozak, J. Mol. Biol. 196, 947 (1987).

19

Conclusions

fragment, which may be suitable for structural investigations by means of In vitro selection of random rRNA fragments (SERF) is a simple and straightforward method to find the native rRNA-binding site for ribosomal proteins. As demonstrated, it has the potential to find a minimal rRNA

The atomic structure of 15 ribosomal proteins has been solved, and one NMR spectroscopy or X-ray crystallography.

S5, S8, L1, L6, and L9. One possible strategy is to separate the domains of those proteins by genetic means and to perform the selection with each RNA-binding sites. It might be difficult to find both interactions in a single SERF experiment. However, in some cases the proposed binding sites are located on different domains, examples include the ribosomal proteins S4, of the surprising findings was that many of the proteins have two potential domain separately.

Acknowledgments

We thank Dr. François Franceschi for kindly providing purified L11 from Thermus thermophilus, Sean Connell for help and discussions, and Detley Kamp for expert assistance in the purification of E. coli ribosomal proteins.

[19] Optimized Synthesis of RNA-Protein Fusions for in Vitro Protein Selection

By RIHE LIU, JEFFREY E. BARRICK, JACK W. SZOSTAK, and RICHARD W. ROBERTS

Introduction

amplify protein molecules that have been selected for function. In order to isolate peptides or proteins with a desired function, the genetic informaaddress. A major advantage of fully in vitro approaches is that they allow iterated cycle of selection and amplification, there is no simple way to tion must be kept topologically linked to the protein in the form of a coding tems is largely mediated by polypeptides. The difficulty in designing schemes for in vitro protein selection is that while in vitro selection involves an sequence such as RNA or DNA, a set of chemical tags, or a physical of peptides and proteins represents an area of great interest, in large part because the chemistry of molecular recognition and catalysis in living sys-The extension of in vitro selection technology to the in vitro selection

Copyright © 2000 by Academic Press All rights of reproduction in any form reserved.

Oct. 6879/00 \$30.00

RNA-PROTEIN FUSIONS

the isolation or proteins with desired properties even when no in vivo selection strategy exists or can be designed

experiment. For example, RNA molecules that bind ATP occur with a selection designed to isolate ATP-binding aptamers would be unlikely to complexity) is one of the most important variables in a combinatorial frequency of 1/1011 in random sequence RNA libraries. 12 Thus, an RNA identify those individuals that have the desired functional properties. The number of different molecules that can be examined (the pool size or or pool containing many different sequences. This pool is then sieved to In vitro selection experiments begin with the generation of a population

Until recently, in vivo and in vitro protein selection experiments (e.g., succeed if the starting library contained only 1 billion sequences.

amined in 1961,8 has been the subject of previous reports and will not be covered in detail here.9-11 experiments routinely involve generating and screening libraries containing more than 10^{15} independent sequences. Two approaches have been developed that provide for totally in vitro selection of peptides and proteins: ribosome display and mRNA-protein fusions. Ribosome display, first extion on library size results from transfecting the starting cDNA library into the organism of choice. In contrast, in vitro RNA and DNA selection ities of about 1 million to 1 billion molecules, respectively. The main limitathe yeast two-hybrid system³ and phage display⁴⁻⁷) were limited to complex-

cin at their 3' end. Because the coding and polypeptide sequences are mRNA-protein fusions for in vitro protein selection. An mRNA-protein fusion consists of a protein sequence covalently linked via its C terminus to the 3' end of its own messenger RNA (Fig. 1). 12 The fusions are generated by in vitro translation of appropriate mRNA templates containing puromy-This article describes improvements in the development and use of

¹ M. Sassanfar and J. Szostak, Nature **364**, 550 (1993)

² D. H. Burke and L. Gold, Nucleic Acids Res. 25, 2020 (1997).
³ S. Fields and O.-K. Song, Nature 340, 245 (1989).

4 G. P. Smith, Science 228, 1315 (1985).

5 J. K. Scott and G. P. Smith, Science 249, 386 (1990).

J. Devlin, L. C. Panganiban, and P. E. Devlin, Science 249, 404 (1990).
 S. E. Cwirla, E. A. Peters, R. W. Barrett, and W. J. Dower, Proc. Natl. Acad. Sci. U.S.A.

⁸ D. B. Cowie, S. Spiegelman, R. B. Roberts, and J. D. Duerksen, Proc. Natl. Acad. Sci.

⁹L. C. Mattheakis, R. R. Bhatt, and W. J. Dower, Proc. Natl. Acad. Sci. U.S.A. 91, 9022

10 L. C. Mattheakis, J. M. Dias, and W. J. Dower, Methods Enzymol. 267, 195 (1996).

Hanes and A. Pluckthun, Proc. Natl. Acad. Sci. U.S.A. 94, 4937 (1997).
 W. Roberts and J. W. Szostak, Proc. Natl. Acad. Sci. U.S.A. 94, 12297 (1997).





SHEET	Serial Number		
DUPLEX DOCUMENT INDEX SH	371P PCT Papers in a 371P Application FOR Foreign Reference NPL Non-Patent Literature FRPR Foreign Priority Papers		
DUPLEXI	Date	Doc Code	Pages

DUPLEX

Employee ID

PCT

I

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

WO 00/21909 20 April 2000 (20.04.00)

(51) International Patent Classification 7:	42	(11) International Publication Number:
CO/B GD/O	9	(43) International Publication Date:
(21) International Application Number:	PCT/US99/23444	PCT/US99/23444 (81) Designated States: AE, AL, AM, A
(22) International Filing Date: 7	7 October 1999 (07.10.99)	ES, FI, GB, GD, GE, GH, GM, P

9 October 1998 (09.10.98) (30) Priority Data: 09/169,426

S

(71) Applicant: PHARMACOPEIA, INC. [US/US]; 3000 Eastpark Boulevard, Cranbury, NJ 08512 (US).

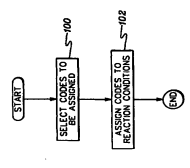
(72) Inventors: DILLARD Lawrence, Wayne; 278 Springhill Road, Skillman, NJ 08558 (US). CONNELLY, James, Andrew; 1865 W. Desert Porat, Oov Valley, AZ 85737 (US). BALDWIN, John, J.; 621 Gypay Hill Circle, Gwynedd Valley, PA 19437 (US). HORLBECK, Eric, George: 17784 Calle de la Siena, San Diego, CA 92130 (US). KIRK, Gregory, L.; 23 Jefferson Road, Winchester, MA 01890 (US). LAURI, Giorgoc, 8 Arthurs Round Table, Wynnewood, PA 19096 (US).

(74) Agent: SCHILLER, Blanche, E.; Heslin & Rothenberg, P.C., 5 Columbia Circle, Albany, NY 12203 (US).

Designated States: AE, AI, AM, AT, AU, AZ, BA, BB, BG, BB, BY, CA, CH, CH, CG, CG, CG, DB, DM, DM, EE, BF, FI, GB, GD, GB, GH, GM, RB, HU, DI, LI, NI, S. PF, KE, KG, KC, KE, LK, LB, LS, LT, LU, LV, MD, MG, MK, NM, MW, NG, NG, RZ, PF, RO, RU, SD, SE, SG, SI, SK, ST, TI, TM, TR, TT, UA, UG, UZ, VM, YU, ZA, ZW, ARPO patent (GH, GM, KE, LS, MW, SS, SL, SZ, TZ, UG, ZW), Eurasin spatent (AM, AZ, BY, KG, KZ, MD, RU, TI, TM), European generi (AT, BE, CH, CY, DE, DK, ER, SF, FR, RG, GR, ME, TI, LU, MC, NL, PT, SB, OAP) patent (GF, BI, CC, CI, CM, GA, GN, ML, MR, NE, SN, TD, TO).

Published Without international search report and to be republished upon receipt of that report.

(54) Title: SELECTING CODES TO BE USED FOR ENCODING COMBINATORIAL LIBRARIES



(57) Abstract

Codes to be used for encoding combinatorial libraries are selectively chosen based on one or more predefined function or criterion. In particular, a subset of N possible codes is selected based on some criterion. In one example, the codes are binary codes, and each code represents the tags used during a particular stage of synthesis of members of a combinatorial library. The tags define the reaction condition tag, that particular stage of synthesis. In one embodiment, the predefined criterion ensures that each code includes more than one tag. This helps eliminate ambiguity during a decoding process in which the tags are identified to determine the reaction history during synthesis.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

	Slovenia					TG Togo																			
	8	22	83	S.	F	۵	F	F	F	۲	3	-	-	>	>	~	2								
	Lesotho	Lithuania	Luxembourg	Lervia	Monaco	Republic of Moldova	Madagascar	The former Yugoslav	Republic of Macedonia	Mali	Mongolia	Mauritania	Malawi	Mexico	Niger	Netherlands	Norway	New Zealand	Poland	Portugal	Romania	Russian Federation	Sudan	Sweden	Singapore
	3	5	3	2	Ř	æ	¥C	X		Æ	X	AR	ž	XX	ž	Z,	Ş	ž	굼	F	2	2	B	S	S
	Spain	Finland	Prance	Gebon	United Kingdom	Georgia	Ghana	Guinea	Greece	Hungary	Ireland	Israel	Iceland	Italy	Japan	Kenya	Kyrgyzsten	Democratic People's	Republic of Korea	Republic of Korea	Kazakstan	Saint Lucia	Liechtenstein	Sri Lenka	Liberia
	ន្ទ	E	æ	Ş	89	GE	ij	Š	ğ	2	Ħ	=	S	E	4	KE	КĞ	ğ		Ä	Ž	3	3	ž	3
	Albania	Armenia	Austria	Australia	Azerbaijan	Bosnia and Herzegovina	Barbados	Betgium	Burkina Faso	Bulearia	Benin	Brazil	Belans	Cenada	Central African Republic	Congo	Switzerland	Côte d'Ivoire	Сатегооп	China	Cubs	Czech Republic	Germany	Denmark	Batonia
	¥.	¥	1	7	77	ВА	88	BE	BF	8	2	#	ВУ	ర	ð	8	ö	ប	3	3	5	ដ	DE	ΩĶ	E
-		_	_			-	_	_	_	_			_		_								_	_	

SELECTING CODES TO BE USED FOR ENCODING COMBINATORIAL LIBRARIES

TECHNICAL FIELD

This invention relates, in general, to the encoding of combinatorial libraries and, in particular, to selectively choosing codes to be assigned to reaction conditions used during synthesis of a combinatorial library.

BACKGROUND ART

challenge in using combinatorial libraries is the characterization of members of the temperature changes, etc.) for each step. The complexity, or number of members conditions for each step of the synthesis, and can therefore, be quite large. The molecular libraries having immense diversity. These techniques entail a series of Combinatorial techniques of chemical synthesis allow the creation of in a combinatorial library, is given by the product of the number of reaction chemical steps with multiple choices of reaction conditions (e.g., reagents,

2

library with particular desired properties. 15

Through a protocol of separating and mixing beads during the synthesis, each bead divide technique to perform chemical synthesis on solid particles, such as beads. chemically bound to it, and that product is likely to differ from that bound to One solution to the above challenge is to use a split synthesis or direct in the final library has a product from a single, specific reaction sequence 20

another bead.

molecules) are attached to each bead in order to encode the reaction condition During each step of the synthesis, zero or more tags (e.g., tagging used in that step, as well as the step number.

WO 00/21909

PCT/US99/23444

111 (Reagent 7). Thus, a binary synthesis code describing any complete N-STEP combinatorial synthesis using any of seven different reagents in each of N steps is products. As an example, the various reagents which can be used in any step are In one embodiment, the tags are used in combination with one another to form a binary record of the synthetic steps for each bead. For example, assume a to be carried out. Such a combinatorial synthesis would yield $7^{\rm N}$ different final designated as binary 001 (Reagent 1), 010 (Reagent 2), 011 (Reagent 3)... synthesis using $3 \times N$ binary digits can be written.

This 9-bit binary synthesis code describes the synthesis, and can be read from right 011. Further, if Reagent 6 is used in the third step, the description is 110 001 011. description is 011. If Reagent 1 is used in the second step, the description is 001 to left in 3-bit blocks to decode the reagents used in each step of the synthesis. For instance, if Reagent 3 is used in the first step, the binary numerical 2

molecules, T9-T1, for the above example, where T9 represents the leftmost binary tag represents a binary "1" for the corresponding bit. Using a set of nine tagging bit and T1 represents the rightmost bit, the tag mixture containing only T9, T8, sensitively detectable molecules is used as tags, and the presence of a particular To represent such a synthesis code chemically, a set of distinguishable, T4, T2 and T1 represents the 110 001 011 synthesis code. 15

- libraries indexed with molecular tags", Proceedings Of The National Academy Of The use of tags and various encoding techniques are described in detail in herein by reference in its entirety. Ohlmeyer et al., "Complex synthetic chemical Sciences Of The United States Of America, Vol. 90, No. 23, pp. 10922-10926 one or more of the following references, each of which is hereby incorporated 20
 - (December 1993); J.J. Baldwin, "Design, synthesis and use of binary encoded synthetic chemical libraries", Molecular Diversity, Vol. 2, No. 1/2, pp. 81-88 encoded combinatorial libraries", Proceedings Of The National Academy Of (October 1996); Burbaum et al., "A paradigm for drug discovery employing 52

Sciences Of The United States Of America, Vol. 92, No. 13, pp. 6027-6031 (June Chemical Libraries Encoded With Tags", issued on October 15, 1996; Baldwin et 1995); Still et al., U.S. Patent No. 5,565,324, entitled "Complex Combinatorial al., U.S. Patent No. 5,618,825, entitled "Combinatorial Sulfonamide Library",

- 'Synthesis Of Combinatorial Libraries", issued on September 02, 1997; Still et al., Chemical Libraries Encoded With Tags", International Publication Date April 14, international Publication No. WO 94/08051, entitled "Complex Combinatorial issued on April 08, 1997; Baldwin et al., U.S. Patent No. 5,663,046, entitled 1994; Dower et al., U.S. Patent No. 5,639,603, entitled "Synthesizing And
- Diverse Collections Of Oligomers", International Publication Date April 01, 1993. International Publication No. WO 93/06121, entitled "Method Of Synthesizing Screening Molecular Diversity", issued on June 17, 1997; and Dower et al., 2

While binary coding has been established as a viable technique in encoding complex combinatorial libraries, there are some shortcomings with the present

techniques, especially during decoding of the tags. 12

detached and identified to determine the particular conditions that occurred during synthesis. One technique for separating and identifying tags is known as Capillary particular bead. In particular, during decoding, any tag(s) attached to a bead is Decoding is performed in order to determine the reaction history of a Gas Chromatography (GC)

8

present due to impurities of similar retention times or tags present in low amounts. During decoding, it is sometimes difficult to determine whether a tag is circumstances, to determine the appropriate binary code that represents the Thus, the decoding becomes ambiguous, and it is difficult, under those reaction history.

eliminates ambiguous code reading. A further need exists for a capability that Based on the foregoing, a need exists for a coding technique that

22

enables the selective choosing of codes, from N possible codes, to be assigned to reaction conditions. A yet further need exists for a capability that guarantees the significant variability in the absolute timing of the run, the relative timing can be presence of enough tag peaks in a chromatogram that, even in the presence of

determined, even if there is significant variability in their positions (i.e., absolute determined. That is, a need exists for the presence of at least two tag peaks, so timing). A further need exists for a capability that uses this time distance to that the time distance (i.e., relative timing) between those peaks can be confirm whether a particular peak represents the presence of a tag or a

tag). The use of time spacing as a confirmation of whether a peak represents a tag contaminant (e.g., substantially constant time spacing indicates the presence of a or a contaminant is referred to herein as "self-clocking". 2

SUMMARY OF THE INVENTION

plurality of codes to be assigned to a plurality of reaction conditions usable during The shortcomings of the prior art are overcome and additional advantages plurality of tags, wherein none of the plurality of codes includes only a single tag. encoding combinatorial libraries. The method includes, for instance, selecting a are provided through the provision of a method of determining codes usable in synthesis of a combinatorial library. Each of the plurality of codes includes a The method further includes assigning selected codes to reaction conditions.

15

In one embodiment, each of the plurality of codes is a binary code, and a binary "one" within the binary code represents the presence of a particular tag.

ឧ

using a predefined criterion. The predefined criterion specifies at least one of the In another embodiment of the invention, the plurality of codes is selected following: each of the plurality of codes includes an even number of tags present therein; each of the plurality of codes includes up to a maximal number of tags therein; each of the plurality of codes includes an odd number of tags present 23

In another embodiment, the plurality of codes is selected using a parity bit.

In a further aspect of the present invention, a method of determining codes usable in encoding chemical libraries is provided. The method includes selecting, from N possible codes, a group of codes to be assigned to a plurality of reaction using a predefined function to select the group of codes, wherein the predefined conditions usable during synthesis of a chemical library. The selecting includes function selects fewer than N-1 codes from the N possible codes. The method further includes assigning selected codes to reaction conditions. 2

selecting, from N possible codes, a plurality of codes to be assigned to a plurality In yet a further aspect of the present invention, a method of determining criterion. The predefined criterion is other than excluding an "all zeroes" code. codes usable in encoding chemical libraries is provided. The method includes The method further includes assigning selected codes to reaction conditions. of reaction conditions, wherein the plurality of codes satisfies a predefined

13

conditions, in which each of the plurality of codes includes a plurality of tags, such usable in encoding combinatorial libraries is provided. The system includes means In a further aspect of the present invention, a system of determining codes that none of the plurality of codes includes only a single tag. The system further for selecting a plurality of codes to be assigned to a plurality of reaction includes means for assigning selected codes to reaction conditions.

ន

selecting, from N possible codes, a group of codes to be assigned to a pluraity of usable in encoding chemical libraries is provided. The system includes means for In another aspect of the present invention, a system of determining codes 22

WO 00/21909

PCT/US99/23444

possible codes. The system further includes means for assigning selected codes to codes, wherein the predefined function selects fewer than N-1 codes from the $\ensuremath{\mathrm{N}}$ reaction conditions usable during synthesis of a chemical library. The means for selecting includes means for using a predefined function to select the group of

reaction conditions.

means for selecting, from N possible codes, a plurality of codes to be assigned to a In yet a further aspect of the present invention, a system of determining codes usable in encoding chemical libraries is provided. The system includes

"all zeroes" code. The system further includes means for assigning selected codes predefined criterion, wherein the predefined criterion is other than excluding an plurality of reaction conditions, wherein the plurality of codes satisfies a to reaction conditions. 2

In another aspect of the present invention, an article of manufacture is

includes only a single tag, and computer readable program code means for causing codes usable in encoding combinatorial libraries. The computer readable program conditions usable during synthesis of a combinatorial library, each of the plurality readable program code means embodied therein for causing the determining of computer to select a plurality of codes to be assigned to a plurality of reaction of codes including a plurality of tags, wherein none of the plurality of codes code means includes computer readable program code means for causing a provided, including at least one computer usable medium having computer a computer to assign selected codes to reaction conditions. 2 15

selecting, from N possible codes, a group of codes to be assigned to a plurality of In yet another aspect of the present invention, at least one program storage one program of instructions executable by the machine to perform a method of reaction conditions usable during synthesis of a chemical library. The selecting device is provided, which is readable by a machine, tangibly embodying at least determining codes usable in encoding chemical libraries. The method includes 23

WO 00/21909

predefined function selects fewer than N-1 codes from the N possible codes. The includes using a predefined function to select the group of codes, wherein the method further includes assigning selected codes to reaction conditions.

selecting, from N possible codes, a plurality of codes to be assigned to a plurality In a further aspect of the present invention, at least one program storage one program of instructions executable by the machine to perform a method of criterion. The predefined criterion is other than excluding an "all zeroes" code. device is provided, which is readable by a machine, tangibly embodying at least determining codes usable in encoding chemical libraries. The method includes The method further includes assigning selected codes to reaction conditions. of reaction conditions, wherein the plurality of codes satisfies a predefined

2

the capability of the present invention guarantees the presence of enough tag peaks absolute timing of the run, the relative timing can be determined. Additionally, the invention in each synthesis step can result in the avoidance of single bit codes for capability of the present invention advantageously enables the selective choosing of codes, from N possible codes, to be assigned to reaction conditions. Further, in a chromatogram that, even in the presence of significant variability in the capability is provided that eliminates ambiguous code reading. Further, the In accordance with the principles of the present invention, a coding present invention is advantageously self-clocking. The use of the present reaction conditions. 8

12

Additional features and advantages are realized through the techniques of described in detail herein and are considered a part of the claimed invention. the present invention. Other embodiments and aspects of the invention are

25

BRIEF DESCRIPTION OF THE DRAWINGS

WO 00/21909

PCT/US99/23444

φ

specification. The foregoing and other objects, features, and advantages of the The subject matter which is regarded as the invention is particularly invention will be apparent from the following detailed description taken in pointed out and distinctly claimed in the claims at the conclusion of the conjunction with the accompanying drawings in which:

FIG. 1 depicts a generalized method of the present invention;

~

FIG. 2 depicts one embodiment of the logic associated with the selection capability of the present invention;

is selected (or "accepted") therefrom, in accordance with the principles of FIG. 3 depicts one example of N possible codes, in which a subset the present invention;

10

FIG. 4 depicts one example of a table of accepted codes, in accordance with the principles of the present invention; FIG. 5 depicts one embodiment of the table of accepted codes of FIG. 4, in which the codes are assigned to reaction conditions, in accordance with the principles of the present invention;

15

FIG. 6 depicts another example of accepted codes, in accordance with the principles of the present invention;

FIG. 7a depicts another example of N possible codes, in which a subset is selected (or accepted) therefrom, in accordance with the principles of the present invention;

8

possible codes of FIG. 7a in order to make a selection of the codes to be FIG. 7b depicts one example of adding a parity bit to the N

٠ţ

included in the subset of accepted codes, in accordance with the principles of the present invention;

FIG. 8 depicts one example in which it is difficult to determine whether a tag is present in a sample; FIG. 9 illustrates one example of output from a gas chromatograph, in accordance with the principles of the present invention;

chromatograph, in accordance with the principles of the present invention; FIG. 10 depicts another example of output from a gas and

FIG. 11 depicts one embodiment of a computer environment providing and/or using the capability of the present invention.

2

BEST MODE FOR CARRYING OUT THE INVENTION

chosen based on one or more criterion. That is, out of N possible codes, a subset of the N codes is selected based on some constraint or some predefined function. In accordance with the principles of the present invention, a capability is provided in which codes to be assigned to reaction conditions are selectively

15

The codes are, for instance, binary codes, and each code represents the tags used during a particular stage of synthesis of members of a combinatorial library. Specifically, the tags define the reaction condition used during that

particular stage of synthesis. 2

selectively chosen, from N possible codes, based upon a predefined function or

described with reference to FIG. 1. Initially, a group of acceptable codes is

A generalized technique of one embodiment of the present invention is

WO 00/21909

<u>-</u>10

PCT/US99/23444

arbitrary or may be based on one or more factors. For instance, the first code may selected codes are assigned to reaction conditions, and those conditions may be one or more criterion, as described in detail below, STEP 100. Thereafter, the used during synthesis of library members, STEP 102. The assignment may be

be assigned to the first condition to be used during synthesis, etc.

for a particular synthesis step, STEP 200. For example, a determination is made One embodiment of the selection process used to choose a group of codes FIG. 2. Initially, a decision is made as to the N possible codes that could be used to be assigned to the reaction conditions is described in detail with reference to

as to how many bits are sufficient to represent the number of reaction conditions sufficient. Thus, in a binary scheme, there are 2^4 possible codes (i.e., N=16), as to be used during that step. Assume for this one example that four bits are shown in FIG. 3. 2

parity; an odd parity, codes that do not include a predefined sequence of bits (e.g., the codes that do not include a sequence of 101), etc. The above criteria are only an odd number, greater than one, of "one" bits; more than one tag (i.e., more than one binary "one" bit); up to a total number of "zero" bits; up to a total number of different possibilities. As examples, the criterion can select codes having an even number of "one" bits (i.e., an even number of tags); an odd number of "one" bits, "one" bits; up to a maximal allowed sequence of "zero" or "one" bits; an even criterion (or predefined function) are selected. The criterion can include many Out of the possible codes, only those codes that satisfy one or more 20 15

includes all those codes that have up to a total of two "zeroes". Thus, the process For illustration purposes only, the predefined function, in this example, continues with determining which of the codes of FIG. 3 meet this criterion. 23

provided as examples. There are many more possibilities, and each of those

possibilities is considered a part of the claimed invention.

Returning to FIG. 2, a code is considered from the possible codes, STEP 202. For example, code 0000 is considered from the codes depicted in FIG. 3. A determination is made as to whether the considered code meets the criterion of having no more than two zeroes, STEP 204. Since code 0000 has more than two zeroes, it does not meet the criterion, and thus, is not an acceptable code. Therefore, if there are further codes to be considered, INQUIRY 206, a new code is considered, STEP 202.

Proceeding sequentially down the list of the possible codes illustrated in FIG. 3, the next code is 0001. Again, since code 0001 does not meet the criterion, STEP 204, and there are more codes, INQUIRY 206, another code is selected. This procedure continues. At some point, a code is selected that does meet the criterion, INQUIRY 204. For instance, code 0011 meets the criterion of having at a maximum two "zeroes", thus, this code is selected as an acceptable code.

2

In one example, the acceptable code is placed in a table of acceptable codes, STEP 208. One instance of such a table is depicted in FIG. 4. The above process continues until all of the acceptable codes are selected from the possible codes based on the predefined criterion.

15

As described with reference to FIG. 1, after the acceptable codes are determined, each of the acceptable codes can be assigned, respectively, to one of a plurality of reaction conditions, as depicted in FIG. 5. In this particular case, ten reaction conditions are represented by ten unique codes.

ន

Another example of a predefined function used to select acceptable codes from N possible codes is one in which all codes having an even number of tags is selected. This is depicted in FIG. 6. In this particular example, five bits are

25 considered necessary to represent the various reaction conditions to be used during synthesis. Thus, there are 32 possible codes.

-12-

PCT/US99/23444

15 codes are acceptable (designated by "used" in the table). That is, 15 codes meet the criterion of having an even number of tags.

In yet another embodiment of the present invention, parity bits may be added to the N possible codes in order to determine the acceptable codes. For example, assume there are eight possible codes, as shown in FIG. 7a. Further, assume that even parity is to be used, in this example. Thus, for each code in which the addition of a binary "one" provides an even number of "one" bits, a parity "one" bit is added, as shown in FIG. 7b.

Thereafter, the codes having the parity "one" bit are selected from the N possible codes. In the example depicted in FIG. 7b, the following codes are chosen: 0011, 0101, 1001 and 1111.

2

In the above examples in which each code is a binary code, each binary "one" represents the existence of a tag and each binary "zero" represents the absence of a tag. When a reaction condition is used in a particular synthesis step,

15 the tags represented by the binary code associated with that reaction condition are also added during that synthesis step. This provides a record of the conditions used during synthesis of a particular library member. For example, assume Reaction Condition 3 (FIG. 5) is used during a first synthesis step, Reaction Condition 1 is used during a second step, and Reaction Condition 6 is used during a third step, then the synthesis code representing the three reaction conditions is 0111 0011 0110. This synthesis code can be read from right to left, in 4-bit blocks, to decode the reaction conditions used during each step of the synthesis. (In another embodiment, the 10 different reaction conditions in FIG. 5 would be alternatives for a single reaction step. A different set of

2

25 reaction conditions, encoded by a different set of tags, would be used for the next reaction step.)

this specific example, twelve (12) bits were used to represent 3 reaction conditions To represent the synthesis code chemically, a set of distinguishable tags is used, in which the presence of a particular tag is represented by a binary "1". In and 3 steps and, thus, a set of twelve tagging molecules are used, T12-T1. T12 synthesis code. Therefore, the following tags would represent the 0111 0011 represents the leftmost binary bit and T1 represents the rightmost bit of the 0110 synthesis code: T11, T10, T9, T6, T5, T3 and T2.

which tagging molecules relate to the binary bits of a synthesis code, are known in the art and described in various of the above references, each of which has been The manner in which tagging molecules are prepared and the manner in incorporated herein by reference in its entirety.

2

produced showing peaks where tags are present. From the peaks, a binary code is show that T2 and T3 are present and T1 and T4 are absent, the binary code 0110 is provided, This code indicates that Reaction Condition 3 was used during the determined. For instance, in the above example, since a chromatogram would associated with that bead are decoded. In one example, a chromatogram is When the reaction history of a particular bead is desired, the tags first synthesis step. 15

similar to that of one of the tags in the set. Both of these occurrences can result in chromatogram data in which the tagging code is ambiguous. An example of such impurities can be introduced into the tag mixture, which may have retention times it is difficult to determine whether a tag is present. For instance, it is possible that during the tagging reaction, one tag may not be incorporated in a particular bead Sometimes the peaks of the chromatogram are not well defined, and thus, at the same quantitative level as others within the set. It is also possible that during the detagging/gas chromatograph (GC) portion of the process, some is depicted in FIG. 8. 25

2

WO 00/21909

<u>-</u>

tags 5 and 2 are present, while tags 1 and 3 are not. However, because the peak having the retention time in the expected range for tag 4 is of much lower height In the example of FIG. 8, a 5 place binary code is represented. Clearly, than either of the other two, its identity is in question. As a result, the binary

code, from left to right, can be either 11010 or 10010.

from the 32 (2^3) (see FIG. 6) possible codes, based on a selection criterion of only those codes where the sum of the individual digits is even, then 11010 would not be an acceptable code. Thus, 10010 must be the code that represents the sample possible binary codes is employed. For example, if a group of codes is selected depicted in FIG. 8. With this strategy, $2^{(k+1)} \cdot 1$ unique sets of conditions can be $T_{\rm O}$ eliminate this type of ambiguous code reading, the encoding protocol of the present invention is used, in which a specially chosen subset of the $\ensuremath{\mathsf{N}}$ 2

chromatogram, and each tag within the chromatogram is labelled by CnnCLn. For invention is used are depicted in FIGs. 9-10. Each of the figures depicts a sample Further examples in which the selective code capability of the present example, in FIG. 9, peak 900 has tag C12CL.5. 12

encoded with N tags.

example (e.g., an even number of tags), binary code 1001 (reference number 904) 1101 or 1001. However, based on the criterion for acceptable codes used in this must be the correct code. Therefore, in accordance with the present invention, presence of a particular tag or not. Thus, the binary code for that step may be $\ln {\rm FIG.}~9,$ there is an ambiguity as to whether peak 902 represents the that peak does not have a tag. ន

determined to be tagged. Thus, a binary "one" represents that peak (see reference Similarly, in FIG. 10, peak 1000 is in question. However, based on the present invention in which the criterion specifies no single tags, the peak is number 1002). The same holds true for peak 1004.

-15-

The capability of the present invention can readily be automated by creating a suitable program, in software, hardware, microcode, firmware or any combination thereof. Further, any type of computer or computer environment can be employed to provide, incorporate and/or use the capability of the present invention. One such environment is depicted in FIG. 11 and described in detail helow.

In one embodiment, a computer environment 1100 includes, for instance, at least one central processing unit 1102, a main storage 1104, and one or more input/output devices 1106, each of which is described below.

2

As is known, central processing unit 1102 is the controlling center of computer environment 1100 and provides the sequencing and processing facilities for instruction execution, interruption action, timing functions, initial program loading and other machine related functions. The central processing unit executes at least one operating system, which as known, is used to control the operation of the computing unit by controlling the execution of other programs, controlling communication with peripheral devices and controlling use of the computer resources.

15

Central processing unit 1102 is coupled to main storage 1104, which is directly addressable and provides for high speed processing of data by the central processing unit. Main storage may be either physically integrated with the CPU or constructed in stand alone units.

ន

Main storage 1104 is also coupled to one or more input/output devices 1106. These devices include, for instance, keyboards, communications controllers, teleprocessing devices, printers, magnetic storage media (e.g., tape, disks), direct access storage devices, and sensor based equipment. Data is transferred from main storage 1104 to input/output devices 1106, and from the input/output devices back to main storage.

23

WO 00/21909

PCT/US99/23444

-16-

Described in detail above is an improvement of the coding process of combinatorial libraries, in which group coding is used. Group coding includes selectively choosing a subset of codes, from N possible codes, that meets one or more predetermined criterion. The subset can include codes that are selected

5 based on a parity bit or by some other mechanism.

In one example, the code selection capability of the present invention guarantees the presence of enough tag peaks in the chromatogram that, even in the presence of significant variability in the absolute timing of the run, the relative timing can be determined and the "zero" bits identified reliably. The group coding

10 thus is considered self-clocking.

In one embodiment, the selection capability of the present invention allows more than 2⁽⁰⁺¹⁾-1 possibilities for N bits. Further, no bit can be claimed to be merely extraneous to the code. A bit is considered extraneous to the code when the bit is added to the code just for the sake of adding a bit and that bit can really be ignored when the code is read. With the present invention, these extraneous

15 be ignored when the code is read. With the present invention, these extraneous bits are avoided and thus, more efficient use of tags can be made. The use of the present invention at each synthesis step can result in avoiding single-bit codes for reaction conditions. Additionally, it advantageously allows for more reliable interpretation of an individual chromatogram by using the guaranteed presence of a minimum number of tags to create an "internal standard" for the shifts in that chromatogram. Further, it allows for independent error checking of the validity of the tags at each step based on the frequency of the code

Although the examples described above reference binary coding, the present invention is also applicable to higher order coding or other types of coding. Thus, these are considered a part of the claimed invention.

22

occurrence and the identification of invalid codes.

-17-

media. The media has embodied therein, for instance, computer readable program The article of manufacture can be included as a part of a computer system or sold code means for providing and facilitating the capabilities of the present invention. one or more computer program products) having, for instance, computer usable The present invention can be included in an article of manufacture (e.g., separately.

tangibly embodying at least one program of instructions executable by the machine Additionally, at least one program storage device readable by a machine, to perform the capabilities of the present invention can be provided.

performed in a differing order, or steps may be added, deleted or modified. All of The flow diagrams depicted herein are just exemplary. There may be many variations to these diagrams or the steps (or operations) described therein without departing from the spirit of the invention. For instance, the steps may be these variations are considered a part of the claimed invention. 9

from the spirit of the invention and these are therefore considered to be within the modifications, additions, substitutions and the like can be made without departing Although preferred embodiments have been depicted and described in detail herein, it will be apparent to those skilled in the relevant art that various scope of the invention as defined in the following claims. 15

WO 00/21909

PCT/US99/23444

₽-

CLAIMS

What is claimed is:

A method of determining codes usable in encoding combinatorial

2 libraries, said method comprising:

reaction conditions usable during synthesis of a combinatorial library, each of said plurality of codes comprising a plurality of tags, wherein none of selecting a plurality of codes to be assigned to a plurality of said plurality of codes comprises only a single tag; and

assigning selected codes to reaction conditions.

The method of claim 1, wherein each of said plurality of codes is a 2 binary code, and wherein a binary "one" within said binary code represents the 4

presence of a particular tag.

The method of claim 1, wherein said selecting comprises using a

2 predefined criterion to select said plurality of codes from N possible codes.

WO 00/21909

PCT/US99/23444

-16

The method of claim 3, wherein said predefined criterion specifies

2 at least one of the following:

s each of said plurality of codes includes an even number of tags

present therein;

each of said plurality of codes includes an odd number of tags

present therein, wherein said odd number is greater than one;

each of said plurality of codes includes up to a maximal number of

s tags present therein;

each of said plurality of codes includes up to a maximal number of

10 "zero" bits; and

11 each of said plurality of codes does not include a predetermined

12 pattern of bits.

1 S. The method of claim 1, wherein said selecting comprises using a

2 parity bit to determine which codes of N possible codes are to be selected as said

3 plurality of codes.

WO 00/21909

PCT/US99/23444

-20-

A method of determining codes usable in encoding chemical

2 libraries, said method comprising:

selecting, from N possible codes, a group of codes to be assigned

to a plurality of reaction conditions usable during synthesis of a chemical

library, said selecting comprising using a predefined function to select said

group of codes, wherein said predefined function selects fewer than N-1

codes from said N possible codes; and

assigning selected codes to reaction conditions.

7. The method of claim 6, wherein said predefined function specifies

2 that each code of said group of codes has up to a maximal number of "zero" bits.

8. The method of claim 6, wherein said predefined function specifies

2 that each code of said group of codes has up to a maximal number of tags.

The method of claim 6, wherein said predefined function specifies

that each code of said group of codes has up to a maximal number of "one" bits.

10. The method of claim 6, wherein said predefined function specifies

2 that each code of said group of codes has an even number of tags present.

11. The method of claim 6, wherein said predefined function specifies

2 that each code of said group of codes has an odd number of tags present.

12. The method of claim 11, wherein said predefined function specifies

2 that each code of said group of codes has an odd number, greater than one, of tags

present.

-51-

The method of claim 6, wherein said predefined function comprises

2 not including in said group of codes any code having a predetermined pattern.

The method of claim 6, wherein said predefined function comprises 14.

2 using a parity bit to select said group of codes.

The method of claim 6, wherein each of said plurality of codes is a

2 binary code, and wherein a binary "one" within said binary code represents the

3 presence of a particular tag.

16. A method of determining codes usable in encoding chemical

5 libraries, said method comprising:

selecting, from N possible codes, a plurality of codes to be assigned

library, wherein said plurality of codes satisfies a predefined criterion, said to a plurality of reaction conditions usable during synthesis of a chemical

predefined criterion being other than excluding an "all zeroes" code; and

assigning selected codes to reaction conditions.

2

The method of claim 16, wherein each of said plurality of codes is a

2 binary code, and wherein a binary "one" within said binary code represents the

3 presence of a particular tag.

WO 00/21909

-22-

The method of claim 16, wherein said predefined criterion specifies <u>8</u>

2 at least one of the following:

each of said plurality of codes includes an even number of tags

present therein;

each of said plurality of codes includes an odd number of tags present therein; each of said pluraity of codes includes up to a maximal number of tags present therein;

each of said plurality of codes includes up to a maximal number of

each of said plurality of codes does not include a predetermined "zero" bits; and

10

pattern of bits.

Ξ

The method of claim 16, wherein said selecting comprises using a 12

2 parity bit to determine which codes of N possible codes are to be selected as said

3 plurality of codes.

WO 00/21909

PCT/US99/23444

-23-

- A system of determining codes usable in encoding combinatorial 20
- 2 libraries, said system comprising:
- means for selecting a plurality of codes to be assigned to a plurality
- of reaction conditions usable during synthesis of a combinatorial library,
- each of said plurality of codes comprising a plurality of tags, wherein none
 - of said plurality of codes comprises only a single tag; and
- means for assigning selected codes to reaction conditions.
- The system of claim 20, wherein said means for selecting comprises 7
 - 2 means for using a predefined criterion to select said plurality of codes from N
- possible codes.
- 22. A system of determining codes usable in encoding chemical
- 2 libraries, said system comprising:
- means for selecting, from N possible codes, a group of codes to be
- assigned to a plurality of reaction conditions usable during synthesis of a
 - chemical library, said means for selecting comprising means for using a
- predefined function to select said group of codes, wherein said predefined
 - function selects fewer than N-1 codes from said N possible codes; and
- means for assigning selected codes to reaction conditions.
- that each code of said group of codes has up to a maximal number of "zero" bits.

The system of claim 22, wherein said predefined function specifies

23

- The system of claim 22, wherein said predefined function specifies 2 that each code of said group of codes has up to a maximal number of tags.

WO 00/21909

PCT/US99/23444

-24

- The system of claim 22, wherein said predefined function specifies 25.
- 2 that each code of said group of codes has an even number of tags present.
- The system of claim 22, wherein said predefined function specifies **2**6.
 - that each code of said group of codes has an odd number of tags present. 6
- The system of claim 22, wherein said predefined function comprises
- not including in said group of codes any code having a predetermined pattern.
- The system of claim 22, wherein said predefined function comprises 89
- 2 using a parity bit to select said group of codes.
- A system of determining codes usable in encoding chemical 29.
- 2 libraries, said system comprising:
- means for selecting, from N possible codes, a plurality of codes to
- be assigned to a plurality of reaction conditions usable during synthesis of a
 - chemical library, wherein said plurality of codes satisfies a predefined
 - criterion, said predefined criterion being other than excluding an "all
- zeroes" code; and
- means for assigning selected codes to reaction conditions.

WO 00/21909

<u>-</u>5

The system of claim 29, wherein said predefined criterion specifies 30.

2 at least one of the following:

each of said plurality of codes includes an even number of tags present therein;

each of said plurality of codes includes an odd number of tags

present therein;

each of said plurality of codes includes up to a maximal number of

tags present therein;

each of said plurality of codes includes up to a maximal number of

"zero" bits; and 10 each of said plurality of codes does not include a predetermined Ξ

pattern of bits. 2

31. The system of claim 29, wherein said means for selecting comprises

2 means for using a parity bit to determine which codes of N possible codes are to

3 be selected as said plurality of codes.

WO 00/21909

-56-

An article of manufacture, comprising: 32. at least one computer usable medium having computer readable program

3 code means embodied therein for causing the determining of codes usable in

4 encoding combinatorial libraries, the computer readable program code means in

said article of manufacture comprising:

computer readable program code means for causing a computer to

conditions usable during synthesis of a combinatorial library, each of said select a plurality of codes to be assigned to a plurality of reaction

plurality of codes comprising a plurality of tags, wherein none of said

plurality of codes comprises only a single tag; and

computer readable program code means for causing a computer to

assign selected codes to reaction conditions.

11

33. At least one program storage device readable by a machine,

2 tangibly embodying at least one program of instructions executable by the machine

3 to perform a method of determining codes usable in encoding chemical libraries,

said method comprising:

selecting, from N possible codes, a group of codes to be assigned

to a plurality of reaction conditions usable during synthesis of a chemical

library, said selecting comprising using a predefined function to select said

group of codes, wherein said predefined function selects fewer than N-1

codes from said N possible codes; and

assigning selected codes to reaction conditions.

2

PCT/US99/23444

WO 00/21909

PCT/US99/23444

-27-

34. At least one program storage device readable by a machine,

2 tangibly embodying at least one program of instructions executable by the machine

3 to perform a method of determining codes usable in encoding chemical libraries,

4 said method comprising:

selecting, from N possible codes, a plurality of codes to be assigned

to a plurality of reaction conditions usable during synthesis of a chemical

library, wherein said plurality of codes satisfies a predefined criterion, said

predefined criterion being other than excluding an "all zeroes" code; and

assigning selected codes to reaction conditions.

* * * *

SELECT CODES TO
BE ASSIGNED
ASSIGN CODES TO
REACTION CONDITIONS

Fig. 1

3 / 8

PCT/US99/23444

WO 00/21909

ω	
_:	
fig	
4	

TABLE OF ACCEPTED CODES	REACTION CONDITION										
TAE	CODE	0011	1010	0110	1001	1100	0111	1011	1101	1110	1111

DETERMINE N POSSIBLE CODES CONSIDER A CODE FROM NORE V POSSIBLE CODES N Fig. 7 Fig	J1g. 2
--	--------

WO 00/21909

TABLE OF ACCEPTED CODES	REACTION CONDITION	CONDITION 1	CONDITION 2	CONDITION 3	CONDITION 4	CONDITION 5	CONDITION 6	CONDITION 7	CONDITION 8	CONDITION 9	CONDITION 10
TABLE	CODE	0011	0101	0110	1001	1100	0111	1011	1101	1110	1111

NOT USED
NOT USED
USED
NOT USED
USED
NOT USED
USED
NOT USED

